Corrodibility and Adherence of Reinforced Concrete Rebars Electroplated with Zinc and Zinc-Nickel Alloys

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Received: December 19, 2018; Revised: March 11, 2019; Accepted: May 22, 2019

Reinforced concrete is the main structural material used in the world in the construction of bridges, roads and commercial and residential buildings. One effective method for preventing corrosion of steel reinforcement is the application of galvanic coatings to the reinforcement itself. The present study evaluated the performance of three groups of reinforcing steel bars (black steel, used as a reference and rebars electroplated with Zinc and Zinc-Nickel alloys) to corrosion and adherence to the concrete matrix. For the analysis of corrosion, the specimens were subjected to two different accelerated corrosion tests: salt spray exposure and wet and dry semi-cycles. The corrosion potential was measured for the qualitative monitoring of the corrosion process and the corrosion rate was estimated. In order to evaluate the steel-concrete adherence, the pullout test was used, in accordance with RILEM-CEB-FIP. The results showed that Zinc and Zinc-Nickel coatings increased the durability of the reinforcing bars and the accelerated corrosion test of wetting and drying cycles proved to be more aggressive compared to the salt spray method to evaluate the corrosion process in the reinforced concrete specimens. Additionally, galvanized reinforcement reduced adherence to the concrete matrix.

Keywords: electroplating, rebar, corrosion potential, reinforced concrete, adherence.

1. Introduction

Reinforced concrete is the main structural material used in the world in the construction of bridges, roads and commercial and residential buildings. However, corrosive processes in reinforcement bars have resulted in premature deterioration of reinforced concrete structures and generated high costs for maintenance and recovery, especially in countries with urban centers located close to the sea.

According to the U.S. Federal Highway Administration (FHWA), in 2002, the estimated annual direct cost of corrosion in the United States was about $276 billion, approximately 3.1% of the US Gross Domestic Product (GDP). Aging infrastructure (buildings, bridges, roads, plants, pipelines, tanks, and other key elements) is a serious problem with a cost of $22.6 billion 1. In the European Union, 30 to 50% of the annual cost of infrastructure maintenance is spent on corrosion-related issues 2. In Brazil this cost exceeds 3 billion dollars per year 3. This indicates that, in addition to technical issues, the development and use of methods that can maximize the service life of concrete structures is of fundamental economic importance for any country.

Good practices, such as the correct positioning of the reinforcing bars in the moulds and the assuring of correct pouring, good concrete quality, compaction and curing of the concrete structures, would be enough to ease corrosion problems. Knowing the exposure environments is also very important. Additional measures of corrosion protection are available to the engineer and builder 4. These include, but are not limited to:

- the use of membrane-type coatings applied to the surface of concrete
- the painting of concrete
- the impregnation of concrete with materials designed to reduce its permeability
- the addition of corrosion inhibitors to the concrete
- the use of corrosion-resisting materials (e.g. stainless steels or polymeric rebars 5) as a replacement for conventional steel reinforcement
- cathodic protection of the reinforcement; and/or
- the application of galvanic coatings to the reinforcement itself.

The application of galvanic coatings to the reinforcing bars has a special place. Zinc (Zn) is the most frequently used metal in these coatings 6, mainly because of its ability to create a dense and adherent protective film on the bar surface, causing the corrosion rates to decrease compared to those observed in ferrous materials.
In addition to creating a barrier between the steel and the environment, zinc also can galvanically protect the steel. If the coating is damaged and the underlying steel exposed to the environment, zinc (being anodic to iron) will preferentially corrode and sacrificially protect the exposed steel against the corrosion process. In this type of corrosion protection, broadly known as cathodic protection, the base metal becomes the cathode in the corrosion cell and the coating metal the anode. Zinc has a number of characteristics that make it well suited for use as a coating for protecting rebars from corrosion. The excellent field performance of zinc coatings is due to its ability to form dense, adherent corrosion product films and a subsequent rate of corrosion considerably below that of ferrous materials, some 10-100 times slower, depending on the atmosphere.

Zinc is an amphoteric metal, i.e. in solution reacts with both strong acidic and strong bases, the attack being most severe below pH 6 and above pH 13. Between these values, the rate of attack is very slow due to the formation of protective layers on the zinc surface. Zinc in concrete is passivated for pH values between about 8 and 12.5 due to the formation of a protective surface film of corrosion that is relatively insoluble below pH 12.5. Zinc reacts quite vigorously with wet concrete, but this reaction effectively ceases once the concrete has hardened. The result of these reactions is the formation of a barrier layer of calcium hydroxylzincate accompanied by the evolution of hydrogen.

In ordinary concrete, uncoated steel depassivates once the pH level drops below about 11.5, though in chloride-contaminated concrete this depassivation occurs at higher pH levels. In contrast, zinc-coated steel in concrete remains passivated to pH levels of about 9.5 thereby offering substantial protection against the effects of carbonation of concrete. Zinc-coated reinforcement can also withstand exposure to chloride ion concentrations several times higher than that which causes corrosion of rebars.

It is also common to add other components to the Zinc coating, such as cobalt (Co), iron (Fe) and nickel (Ni), forming the alloys Zn-Co, Zn-Fe and Zn-Ni, which present better properties when compared to the pure Zinc coatings. In this context, the compounds based in Zn-Ni are attracting more interest regarding protection against corrosion due to its superior chemical and mechanical properties.

There are two main methods of producing the zinc protective layer: hot-dip galvanizing and electroplating. Hot-dip galvanizing involves the immersion of cleaned steel in a bath of molten zinc at about 450°C allowing a metallurgical reaction to occur between the steel and the zinc. This reaction produces a coating on the steel made up of a series of iron-zinc alloy layers, which grow from the steel/zinc interface with a layer of essentially pure zinc on the outer surface. Electroplating is a process that uses an electric current to reduce dissolved metal cations so that they form a thin coherent metal coating on an electrode. The term is also used for electrical oxidation of anions on to a solid substrate.

Hot-dip galvanizing deposits a thick, robust layer of zinc iron alloys on the surface of a steel rebar. Electroplating produces a layer that is extremely thin and relatively free of pores, avoiding material waste. The thickness of the coating is influenced by some factors, for instance the current density, the salt concentration, the bath temperature, the presence of additives and the nature of the base metal. Furthermore, the electroplating method does not generate intermetallic alloys like the hot dip galvanizing method, providing a more homogenous and thin coating that will not affect mechanic properties of its substrate.

Among the various electrochemical techniques used to monitor the corrosive process in reinforced concrete, corrosion potential measurements are the most commonly used. Monitoring of the corrosion potential enables changes in the electrochemical corrosion process to be recorded, which can be of interest in monitoring structures. The corrosion potential of the rebars is a mixed potential, resulting from the combination of the kinetics of two processes: (1) metal oxidation, and (2) reduction of dissolved oxygen, without providing quantitative information, i.e., information about the reinforcement corrosion rate cannot be obtained. Thus, the rebar corrosion potential provides a rough indication of a rebar corrosion situation or a rebar passive state.

The ASTM C-876/91 standard presents a correlation between corrosion potential ranges and the probability of the occurrence of corrosion as a criterion for corrosion evaluation, taking as reference a copper/copper sulfate electrode (Cu/CuSO₄, Cu²⁺).

Because galvanized steel has a ferritic base which is magnetic, a coating of zinc will not upset the cover. However, it will affect the interpretation of electrochemical test results because the magnitude of the potential will be different between steel and zinc. Once corrosion initiates, the potential of zinc is some 400 mV more negative than that of black (ordinary) steel and therefore it is important to understand how to interpret reference electrode potentials where the criteria suggested in ASTM C876 for the corrosion of black steel simply does not apply. It is important to note that the potential of the zinc and zinc alloys in the coating rises steadily from about -1100 mV (vs SCE) to about -600 mV as the dissolution of the coating proceeds and the more iron-rich alloy layers in the coating are exposed.

According to Sherine et al. and Panek et al., when black steel is electroplated with zinc, lower values of corrosion potential are associated with a high probability of corrosion. These values are equal to -274 mV for ordinary rebars, -953 mV for rebars electroplated with zinc-nickel and -1043 mV for rebars electroplated with zinc, to saturated calomel electrode (SCE), see Table 1.
In reinforced concrete, effective bond between reinforcing bars and concrete is vital for the full development of composite action. Hence, the bond of reinforcing bars plays an important role in the structural behaviour of reinforced concrete. The bond strength between the concrete and reinforcement bar is determined by three main components: shear stresses due to adhesion along the bar surface, the bearing stresses against the faces of rebars (mechanical interlock), and the friction between the rebars and the surrounding concrete. Despite the fact that the principal contribution to bond strength comes from the mechanical interlock, the chemical adhesion mechanism occurs firstly and, after loading, prevents slip that will only happen after the adhesion is destroyed.

The adhesion bond is the result of the chemical reaction between the reinforcement and the hydrated cement compounds within the interstitial transition zone to form a true chemical bond. The nature of the bonding is dependent on the atomic arrangement, molecular conformation, morphological properties and chemical constitution of the reinforcement and concrete and can be attributed to mechanisms including adsorption and wetting electrostatic attraction, chemical bond, reaction bonding, and exchange reaction bonding.

Initially, the corrosion generates an increase in the adherence between rebars and concrete due to the increase in the surface roughness. However, this increase in adhesion occurs only initially because with a higher formation of oxides there is a loss of adherence, reducing the resistant capacity of the concrete. The bond between the concrete and the reinforcement is essential for the full capacity of the reinforcement to be developed and the successful functioning of a reinforced concrete system.

Regarding the galvanized rebars, Cheng et al. argue that due to the low volumetric variation in the zinc corrosion products, there is no significant decrease in the steel-concrete bond during the corrosive process. However, some research indicates that the adherence reduces due to the formation of hydrogen gas when zinc is in contact with fresh concrete. The bond characteristics of electroplated rebars in reinforced concrete have also been studied. Pullout tests were conducted on rebars embedded in the concrete.

### Table 1. Probability of rebars corrosion activity as a function of ranges of corrosion potential for ordinary (black steel) rebars and rebars electroplated with zinc and zinc-nickel alloys (reference: saturated calomel electrode, SCE)

<table>
<thead>
<tr>
<th>Rebar Surface</th>
<th>Probability of Rebar Corrosion Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&quot;Passivation&quot; (&lt; 10%)</td>
</tr>
<tr>
<td></td>
<td>Corrosion (&gt; 90%)</td>
</tr>
<tr>
<td>Black Steel</td>
<td>&gt; -0.124 V</td>
</tr>
<tr>
<td>Electroplated with Zn</td>
<td>&gt; -0.650 V</td>
</tr>
<tr>
<td>Electroplated with Zn-Ni</td>
<td>&gt; -0.550 V</td>
</tr>
<tr>
<td></td>
<td>&lt; -0.274 V</td>
</tr>
<tr>
<td></td>
<td>&lt; -1.043 V</td>
</tr>
<tr>
<td></td>
<td>&lt; -0.953 V</td>
</tr>
</tbody>
</table>

2. Materials and Methods

2.1 Materials

The concrete was produced with Brazilian Portland cement (CPII Z-32, according to the Brazilian NBR 11578 standard), which is equivalent to the ASTM C 596 standard (Pozzolan-modified Portland cement). This cement contains pozzolan and is one of the cements most widely used in the state of Bahia, Brazil. The coarse aggregate was dense crushed granite stone and the fine aggregate was natural siliceous sand commercially available in Salvador, Bahia, Brazil.

Steel bars CA-50 class, according to Brazilian standards, 10.0 mm diameter, were used (C content between 0.3 and 0.5 wt.%), with yield stress greater than 500 MPa and a tensile strength of about 560 MPa. These steel bars are manufactured by the process of hot rolling without subsequent cold work.

2.2 Methods

2.2.1 Characterization of the raw materials and concrete dosage

Physical parameters of the materials such as the specific surface area (estimated by BET, using a Micromeritics Gemini 2370 V1.02 surface area analyzer), particle-size distribution (Mastersizer 2000) and specific gravity (Micromeritics Accupyc 1330 V2.01 helium pycnometer) were determined.

The concrete mix proportion was 1.0 (Portland cement): 1.5 (fine aggregate): 1.3 (coarse aggregate) and the water/cement ratio was 0.5. The mortar content was 75% and the cement consumption was 526 kg/m³. After mixing, a vibrating table was used to ensure efficient compaction.

The corrosion potential was measured using prismatic specimens (50x70x90 mm³) into which steel bars (reference and electroplated) were inserted during molding. For the pull out tests, cylindrical specimens (10 cm in diameter and 10 cm in height) were molded for each type of reinforcement (reference rebars with "surface oxidized", reference rebars without oxidation, zinc coated rebars and zinc nickel coated rebars).

All the specimens were unmolded 24 h after being cast and were cured for 28 days in a humid chamber (> 95% HR). A minimum of 4 samples were tested for each measurement.
2.2.2 Zinc and Zinc Nickel Electroplating

Before the electroplating process, the bars were cleaned with an iron brush and then were immersed in alcoholic aqueous solution to remove impurities and oxides adhering to the surface, assuring the efficiency of electroplating.

The Zn and Zn-Ni alloys were deposited galvanostatically from a deposition solution, with concentrations according to Table 2. Electrodeposition was performed at 25°C in an aerated solution. All the experiments were performed in a single-compartment glass circular cell with a diameter of 7 cm and a capacity of 250 mL. In the deposition experiments, a current source (HP, model 6140A) and a voltmeter-ammeter (ICEL, model ET 208-2B) were used (Figure 1). A bar of graphite, with a diameter of approximately 5 mm was used as an anode, and the steel rebars were used as the substrate for the electrodeposition. The cylindrical steel substrates were placed at 2 cm from the graphite anode in the central region of the cell. The galvanostatic depositions were conducted at a current density of 10 mA/cm² for 8-9 min to produce a 5 µm-thick deposit, which is a typical value in a galvanizing process line.

The mass of the deposit was estimated using the following equation (1)\(^2\):

\[
m_{\text{dep}} = e \cdot S \cdot \rho_D\]

(1)

where \(m_{\text{dep}}\) is the mass of the deposit, “\(e\)” is the thickness of the deposit (5 µm), \(S\) is the deposition surface, and \(\rho_D\) is the density of deposit. The deposit density can be estimated using equations 2 or 3:

\[
\rho_{\text{Zn}} = \rho_{\text{Zn}} \cdot C_{\text{Zn}} = 1.0 \quad (2)
\]

\[
\rho_{\text{Zn-Ni}} = \rho_{\text{Zn}} \cdot C_{\text{Zn}} + \rho_{\text{Ni}} \cdot C_{\text{Ni}} \quad (3)
\]

, where \(\rho_{\text{Zn}}\) is the density of Zn, \(\rho_{\text{Ni}}\) is the density of the Ni, \(C_{\text{Zn}}\) is the Zn mass fraction in the deposit and \(C_{\text{Ni}}\) is the Ni mass fraction in the deposit, estimated from constituents proportion in solution.

According to the microphotographs shown in Figure 2, the obtained coatings had a uniform distribution of electrodeposited ions, creating a homogeneous layer, but with different morphologies for each case. The zinc coating exhibited a surface formed of flat randomly scattered grains of various sizes, and the Zinc-Nickel coating had spherical same sized grains distributed all over the surface, as reported in other research 13,26.

2.2.3 Corrosion potential measurement

The reinforcement steel bars were weighed on an analytical balance with an accuracy of 0.01 g. The area exposed to chloride attack (about 15.83 cm²) was then delimited with electrical insulating tape, as shown in Figure 3. The rebars were positioned so that the exposed area was located in the central region of the specimens, as illustrated in Figure 3d.

There was the possibility of crevice conditions due to the tape. However, after the test, no corrosion was detected on the sites that were covered by tape, indicating that the adhesion between the rebars and the tape was good.

The corrosion potential was verified from a chloride-activated accelerated corrosion test. The electrochemical cell used for corrosion potential measurements was composed of a working electrode, the reinforcement steel rebars and the saturated calomel electrode (SCE) used as the reference electrode.

Before taking the measurements, the side of the specimen to be measured was pre-moistened with a wet sponge for one minute. A conductive solution containing 5 mL of neutral detergent to one liter of water, according to the ASTM C-876/91 standard

<table>
<thead>
<tr>
<th>Component</th>
<th>Zinc Solution (g/L)</th>
<th>Zinc-10%Nickel Solution (g/L)</th>
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</thead>
<tbody>
<tr>
<td>Potassium Chloride</td>
<td>208.0</td>
<td>---</td>
</tr>
<tr>
<td>Zinc Chloride</td>
<td>19.6</td>
<td>38.5</td>
</tr>
<tr>
<td>Nickel Chloride</td>
<td>---</td>
<td>34.5</td>
</tr>
<tr>
<td>Ammonium Chloride</td>
<td>---</td>
<td>150.0</td>
</tr>
<tr>
<td>Boric Acid</td>
<td>20.0</td>
<td>20.0</td>
</tr>
</tbody>
</table>
Corrodibility and Adherence of Reinforced Concrete Rebars Electroplated with Zinc and Zinc-Nickel Alloys

Figure 2. Morphology of film formed on rebar surface after the Zinc (A and B) and Zinc-Nickel (C and D) electroplating process.

Figure 3. (A) and (B) Scheme of delimitation of the bars exposed area; (C) and (D) Position of bars inside the concrete specimen (RIBEIRO [6]).
(Standard Test Method for Half-Cell Potentials of Uncoated Reinforcing Steel in Concrete) and an electrical conductivity of 0.15 ± 0.02 mS/cm were used. For the measurements, the reference electrode was positioned approximately at mid-span over the rebar under analysis. The contact between the reference electrode and the specimen was aided with a damp sponge.

The corrosion test was started after 63 days, when the specimens showed a constant mass (1.0 gram of variation in two consecutive 24-hour readings) and when the measured corrosion potential indicated the formation of a passive film on the surface of the steel rebars \( E_{corr} > -0.124 \) V for reference rebars, \( E_{corr} > -0.650 \) V for zinc coated rebars and \( E_{corr} > -0.550 \) V for zinc-nickel coated rebars). These reference values correspond to a lower than 10% possibility of corrosion occurrence, according to the ASTM C 876/91 standard (for the saturated calomel electrode used in this work).

In other works \(^{27-30}\), it was necessary to define a specific age, or reference age, from which the procedures of the accelerated corrosion tests started. The authors of those studies associated the reference age to the stabilization of the cement hydration process and defined the ages of 63 days \(^{27,28}\) and 80 days \(^{29,30}\) as sufficient for the cement paste to acquire a relatively well developed physical structure and a significantly high degree of hydration.

After reaching the “safe potential,” the specimens were subjected to two different accelerated corrosion tests: i) Salt spray exposure, using a Equilam SS600e chamber in accordance to ASTM B-117 (Standard Practice for Operating Salt Spray (Fog) Apparatus) with NaCl concentration of 5%, pH between 6.5 and 7.2 and temperature of 35°C; and ii) semi-cycles of partial immersion in 3wt% NaCl solution for two days and semi-cycles of drying in a ventilated oven at 50°C for five days. During the semi-cycle of partial immersion, the level of immersion solution was kept at half the height of the specimen. In such conditions, the chloride inflow occurs primarily by capillary absorption, since the specimens are first dried, and after the pores become saturated, the inflow occurs by diffusion, which is accelerated due to water evaporation through the exposed concrete. The corrosion potential \( E_{corr} \) was measured at the end of each semi-cycle.

According to MCCARTER apud SANTOS \(^{24}\), there is a relationship between the saturation level and the capillary suction forces in a porous material. Thus, when there is a dry and exposed region, these suction forces will be greater, pulling the solution faster into the concrete specimen.

At the end of each part of the cycle, the corrosion potential \( E_{corr} \) and the weight of the specimens were measured. The potential measurements were carried out to verify the condition of the rebars regarding its passivation/corrosion state, while the weighing was necessary to verify its saturation level, because if the water absorption after each wet half-cycle was different, the results of corrosion potential would be altered (dry concrete presents high electrical resistivity).

The active or passive state of corrosion was analyzed based on the corrosion potential \( E_{corr} \), using the saturated calomel electrode (SCE) as reference. The test was concluded when two consecutive and full cycles resulted in corrosion potential values below the critical value.

After concluding the test, the rebars were extracted from the samples, cleaned according to the ASTM G-1/03 standard (Standard Practice for Preparing, Cleaning, and Evaluating Corrosion Test Specimens), and weighed to determine weight loss and to compare them with the initial value. Thus, the corrosion rate (CR) can be calculated according to equation (4).

\[
CR = \frac{K \cdot W}{A \cdot T \cdot D}
\]

(4)

where \( K = \text{constant (for CR(\mu m/year)), } K = 8.76 \times 10^7 \); for CR(g/m².year), \( K = 8.76 \times 10^7 \cdot D \); \( W = \text{weight loss (grams)} \); \( A = \text{exposed rebar area (cm²)} \); \( T = \text{exposure time (hours)} \); \( D = \text{steel rebar density (for CA-50 steel, } D = 7.85 \text{ g/cm³). } \) In this study, \( A = 15.83 \text{ cm²}. \)

Metallic zinc reacts strongly with an aqueous solution of hydrochloric acid (HCl) when it is concentrated, and when the metal is exposed for long periods of time. The result is the production of a large amount of hydrogen gas and the formation of zinc chloride (ZnCl) in the solution. In the experiment, the zinc did not react because the HCl solution was diluted and the contact time between the solution and the rebars was only 60 seconds.

2.2.4 Pull out test

A photograph of the testing machine and test setup is presented in Figure 4. The samples were cast with the steel rebar of 10 mm in a vertical position. A ratio of 10:1 between concrete specimen diameter and rebar diameter was used. The steel bar was inserted into the concrete with bond length of 50 mm (Figure 4a). The unbonded segments of the pull-out specimen were created by placing a plastic pipe 50 mm in length around the diameter of the steel bar. The setup for the pullout test is shown in Figure 4b, which is in accordance with RILEM-CEB-FIP \(^{31}\).

All pull out tests were carried out using a universal testing machine EMIC with a load capacity of 300 kN. From the pull out failure load measured by the testing machine, the bond strength was calculated as the failure load divided by the surface area of the bonded length of the reinforcing bar, as per the following equation \(^{50}\).

\[
\sigma = \frac{F}{(\pi \cdot d \cdot l)}
\]

(5)

where \( \sigma = \text{bond strength, } F = \text{failure load, } d = \text{diameter of the reinforcing bar, and } l = \text{bonded length of the reinforcing bar.} \)
2.2.5 Scanning electron microscopy

Scanning electron microscopy (SEM) was used to analyse the coatings of zinc and zinc-nickel on the rebars, verifying their microstructure and chemical composition. A Scanning Electron Microscope (SEM) Philips XL-30 and with EDX microprobe was used (tension of 25 KV and work distance 11-13 mm).

3. Results and Discussion

3.1. Materials characterization

The Portland cement used here has a specific surface area of 0.425 m²/g and a specific gravity of 2.98 kg/dm³. The sand has a specific surface area of 0.68 m²/g, a specific gravity of 2.63 kg/dm³, and fineness Modulus of 1.52, classified as fine sand by the Brazilian NBR 7211 standard. The gravel has a specific gravity of 2.79 kg/dm³ and a maximum dimension of 9.5 mm. The particle size distribution is presented in Figure 5.

3.2. Corrosion potential

The corrosion potential of the rebars exposed to semi-cycles of partial immersion in 3 wt% NaCl solution for two days and semi-cycles of drying in a ventilated oven at 50°C for five days was verified after each semi-cycle. Figure 6 presents the results of the corrosion potential measurements.

In the first 63 days, the specimens were not subjected to wetting and drying cycles in NaCl solution until the “safe potential” was reached, and the tests were interrupted when the “unsafe potential” was reached in two consecutive dry state measurements. The values represent the average of four measurements taken for each composition. The error bars were not placed in graphs because they would make the graphs very confusing and difficult to visualize. However, the results were highly reproducible, with a variation of less than 6%.

One of the aspects of the test procedure adopted is that the rebar corrosion potential varied throughout the test, showing more negative or more positive values depending on the semi-cycle to which the specimen was subjected. The most positive corrosion potential values were recorded after the drying cycles because, due to the decreasing amount of electrolyte, the concentration of dissolved substances increased. In fact, according to the Nernst equation, the equilibrium potential increases with the increase in activity, i.e., the increase in the effective concentrations of oxidized substances. Studies by SANTOS and RIBEIRO et al. confirm this behaviour and show an inverse correlation between the corrosion potential and the moisture content of the concrete, indicating that an increase in moisture content implies a decrease in the measured rebar corrosion potential.
In the salt spray exposure test, the parameters cited before (temperature, humidity, saturation level) are kept constant, causing a different behaviour in the corrosion potential measurements, as can be seen in Figure 7.

As suggested by GONZÁLEZ et al. 32, the measures of the potential are not conclusive by themselves due to the variety of factors that affect it, they merely provide preliminary indicatives of the rebars corrosion situation. As can be seen, the corrosion potential of the coated rebars stayed in the uncertainty range ($E_{cor} > -1.043$ V, for rebars coated with Zinc and $E_{cor} > -0.953$ V, for rebars coated with Zinc-Nickel), while the rebars with no coating presented potentials in the high corrosion probability range ($E_{cor} < -0.274$ V). Such behaviour indicates that there is a protective effect on the reinforcing bars regarding their despassivation when they are coated with Zinc or the Zinc-Nickel alloy by electroplating process, which is due to the delay in the change from the passive to the active state of corrosion.

Figure 6. Evolution of the corrosion potential of reference and coated rebars exposed to semi-cycles of partial immersion in 3 wt% NaCl and drying in a ventilated oven.

Figure 7. Evolution of the corrosion potential of reference and coated rebars exposed to salt spray.
3.3 Corrosion rate

The corrosion rate, calculated from the loss of mass observed after accelerated corrosion tests for each group of rebars (coated with Zinc, Zinc-Nickel and uncoated), for specimens exposed to semi-cycles of partial immersion in 3wt% NaCl solution for two days and semi-cycles of drying in a ventilated oven at 50°C for five days and to salt spray, are presented in Figures 8 and 9, respectively. As expected, the galvanized reinforcement had a lower corrosion rate compared to the reference steel rebars.

The formation of this compound occurs at a pH between 12.0 and 12.8 \(^{34}\) and is associated with the humidity and calcium hydroxide present in the concrete.

\[
\text{Zn} + 2\text{H}_2\text{O} \leftrightarrow \text{Zn(OH)}_2 + \text{H}_2 \quad (6)
\]

\[
2\text{Zn(OH)}_2 + -2\text{H}_2\text{O} + \text{Ca(OH)}_2 \rightarrow \text{Ca}[\text{Zn(OH)}_3]_2 \quad (7)
\]

The results obtained in the salt spray exposure tests (Figure 9) showed a lower variability and a difference of one order of magnitude, which indicates that this test is less aggressive than the wet and dry cycles test. The corrosion rate calculated was equal to 22.50 µm/year for uncoated rebars, 15.58 µm/year for zinc coated rebars, and 15.40 µm/year for zinc-nickel coated rebars.
This is because of capillary suction on the concretes submitted to wet and dry cycles that causes the concentration of chlorides inside the concrete to increase in comparison to the samples exposed to the salt spray.

3.4 Steel-concrete adherence

Figure 10 shows the results of the pull-out test. As expected, a reduction in the adhesion of the galvanized reinforcement in comparison to the reference reinforcement (without cover) was observed, due to the reduction in the height of the ribs. In addition, the chemical iteration between the zinc coatings and the hydrated calcium silicates of the cement may result in different ionic bonds. As a result, a reduction in bond stress of 16% was observed with the application of these treatments.

4. Conclusions

From the results of the tests, it can be concluded that:

Galvanization by electroplating is a protective method which increases the corrosion resistance of rebars used in construction.

The Zinc and Zinc-Nickel coatings increased the durability of the reinforcing bars, although the thickness adopted in this research might not have been enough to promote a substantial increase in the life cycle, because this thickness might have interfered in the performance of the coatings.

The accelerated corrosion test of wetting and drying cycles proved to be more aggressive than the "salt spray method" in the evaluation of the corrosion process in the reinforced concrete specimens because of the capillary suction on the concretes submitted to wet and dry cycles that causes the concentration of chlorides inside the concrete to increase faster.

The use of more refined monitoring techniques is necessary to achieve a better understanding of how the coatings of zinc and zinc-nickel alloy work in reinforced concrete, as there is little in the literature about this.

The adherence loss due to the electrodeposition process is relatively small compared to the increase in the corrosion resistance of these reinforcements.

The adopted adherence test satisfactorily indicated a decrease in the concrete-steel bond for coated rebars which can be associated to the reduction in chemical adhesion.

5. Acknowledgements

CNPq - Brazilian National Council for Scientific and Technological Development.
CAPES - Coordination for the Improvement of Higher Education Personnel
FAPESB - Research Support Foundation of the State of Bahia

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